=> d his ful						
	(FILE 'HOME' ENTERED AT 08:05:56 ON 24 NOV 2009)					
	FILE 'LREGISTRY' ENTERED AT 08:06:34 ON 24 NOV 2009					
	FILE 'REGISTRY' ENTERED AT 08:08:06 ON 24 NOV 2009 ACT HOD829E/A					
L1 L2 L3 L4	SCR 2040 STR STR 1207 SEA SSS FUL L3 AND L2 AND L1					
L5 L6	FILE 'LREGISTRY' ENTERED AT 08:09:02 ON 24 NOV 2009 STR L2 STR L3	9				
L7	FILE 'REGISTRY' ENTERED AT 08:14:22 ON 24 NOV 2009 50 SEA SUB=L4 SSS SAM L5 AND L6					
L8	FILE 'LREGISTRY' ENTERED AT 08:14:51 ON 24 NOV 2009 STR L5	9				
L9	FILE 'REGISTRY' ENTERED AT 08:18:48 ON 24 NOV 2009 3 SEA SUB=L4 SSS SAM L8 AND L6 D SCA					
L10	59 SEA SUB=L4 SSS FUL L8 AND L6					
L11	FILE 'HCAPLUS' ENTERED AT 08:19:43 ON 24 NOV 2009 30 SEA SPE=ON ABB=ON PLU=ON L10					
L12 L13	FILE 'REGISTRY' ENTERED AT 08:21:36 ON 24 NOV 2009 50 SEA SUB=L4 SSS SAM L5 AND L6 1200 SEA SUB=L4 SSS FUL L5 AND L6					
L14	FILE 'HCAPLUS' ENTERED AT 08:22:02 ON 24 NOV 2009 1116 SEA SPE=ON ABB=ON PLU=ON L13					

FILE 'ZCAPLUS' ENTERED AT 08:22:11 ON 24 NOV 2009

FILE 'HCAPLUS' ENTERED AT 08:22:32 ON 24 NOV 2009

QUE SPE=ON ABB=ON PLU=ON ELECTROLYT?

L15

L16 0 SEA SPE=ON ABB=ON PLU=ON L14 (L) L15

FILE 'ZCAPLUS' ENTERED AT 08:23:20 ON 24 NOV 2009

L17 QUE SPE=ON ABB=ON PLU=ON BATTERY# OR BATTERIES#

FILE 'HCAPLUS' ENTERED AT 08:23:32 ON 24 NOV 2009

L18 0 SEA SPE=ON ABB=ON PLU=ON L14 (L) L17

L19 8 SEA SPE=ON ABB=ON PLU=ON L14 AND L15

L20 0 SEA SPE=ON ABB=ON PLU=ON L14 AND L17

FILE 'ZCAPLUS' ENTERED AT 08:25:05 ON 24 NOV 2009

FILE HOME

FILE LREGISTRY

LREGISTRY IS A STATIC LEARNING FILE

CAS INFORMATION USE POLICIES, ENTER HELP USAGETERMS FOR DETAILS.

FILE REGISTRY

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 22 NOV 2009 HIGHEST RN 1193309-59-9 DICTIONARY FILE UPDATES: 22 NOV 2009 HIGHEST RN 1193309-59-9

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH June 26, 2009.

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For informatio on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

FILE HCAPLUS

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after Decembe 26, 1996), unless otherwise indicated in the original publications.

The CA Lexicon is the copyrighted intellectual property of the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or stor of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 24 Nov 2009 VOL 151 ISS 22
FILE LAST UPDATED: 22 Nov 2009 (20091122/ED)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Aug 2009
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

HCAplus now includes complete International Patent Classification (I reclassification data for the third quarter of 2009.

CAS Information Use Policies apply and are available at:

http://www.cas.org/legal/infopolicy.html

This file contains CAS Registry Numbers for easy and accurate substance identification.

During November, try the new LSUS format of legal status information in the CA/CAplus family databases for free! Complete details on the number of free displays and other databases participating in this offer appear in NEWS 10.

FILE ZCAPLUS

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after Decembe 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or stor of this information, without the prior written consent of CAS is strictly prohibited.

FILE COVERS 1907 - 24 Nov 2009 VOL 151 ISS 22
FILE LAST UPDATED: 22 Nov 2009 (20091122/ED)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Aug 2009
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

ZCAplus now includes complete International Patent Classification (I reclassification data for the third quarter of 2009.

CAS Information Use Policies apply and are available at:

http://www.cas.org/legal/infopolicy.html

This file contains CAS Registry Numbers for easy and accurate substance identification.

During November, try the new LSUS format of legal status information in the CA/CAplus family databases for free! Complete details on the number of free displays and other databases participating in this offer appear in NEWS 10.

=> d que 14

SCR 2040 L1

L2STR

1 A -

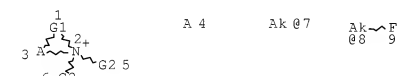
NODE ATTRIBUTES:

CHARGE IS *- AT NSPEC IS RC AT DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 1

STEREO ATTRIBUTES: NONE L3 STR



REP G1 = (2-3) A

VAR G2 = 7/8

NODE ATTRIBUTES:

CHARGE IS *+ ΑT NSPEC IS R ΑT CONNECT IS E1 RC AT DEFAULT MLEVEL IS ATOM

GGCAT IS LOC AT 7
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 9

STEREO ATTRIBUTES: NONE

L4 1207 SEA FILE=REGISTRY SSS FUL L3 AND L2 AND L1

=> d que stat 110

L1 SCR 2040

L2 STR

1 A -

NODE ATTRIBUTES:

CHARGE IS *- AT 1
NSPEC IS RC AT 1
DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

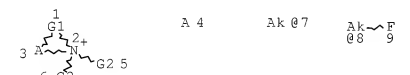
GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 1

STEREO ATTRIBUTES: NONE

L3 STR



REP G1 = (2-3) A

VAR G2=7/8

NODE ATTRIBUTES:

CHARGE IS *+ AT 2
NSPEC IS R AT 4
CONNECT IS E1 RC AT 7
DEFAULT MLEVEL IS ATOM

GGCAT IS LOC AT 7 DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 9

STEREO ATTRIBUTES: NONE

L4 1207 SEA FILE=REGISTRY SSS FUL L3 AND L2 AND L1

L6

A 4 Ak 07 Ak ~ F 08 9

REP G1 = (2-3) A

VAR G2=7/8

VAR G3=C/O/P/S/N

NODE ATTRIBUTES:

CHARGE IS *+ AT 2 NSPEC IS R AT 4

CONNECT IS E1 RC AT 7

DEFAULT MLEVEL IS ATOM

GGCAT IS LOC AT 7

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 9

STEREO ATTRIBUTES: NONE

L8 STR

G1 4

A~F³ @1 F -

VAR G1=2/1

NODE ATTRIBUTES:

CHARGE IS *- AT 1

CHARGE IS *- AT 3

DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 4

STEREO ATTRIBUTES: NONE

L10 59 SEA FILE=REGISTRY SUB=L4 SSS FUL L8 AND L6

100.0% PROCESSED 161 ITERATIONS 59 ANSWERS

SEARCH TIME: 00.00.01

=> d l11 1-30 bib abs hitstr hitind
YOU HAVE REQUESTED DATA FROM FILE 'HCAPLUS' - CONTINUE? (Y)/N:y

L11 ANSWER 1 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 2006:46803 HCAPLUS Full-text

DN 144:135233

TI Pharmaceuticals for inhalation comprising PDE IV inhibitors and glycopyrrolate salts

PA Boehringer Ingelheim Pharma Gm.b.H. & Co. K.-G., Germany

SO Eur. Pat. Appl., 24 pp.

CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
РΤ	 FD 1616567	а 1	20060118	FP 2004-16878	

PI EP 1616567 A1 20060118 EP 2004-16878

200407

16

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU,

PL, SK, HR

CA 2570433 A1 20060126 CA 2005-2570433

200506

```
13
     WO 2006008213
                         Α1
                                20060126
                                           WO 2005-EP52704
                                                                    200506
                                                                    13
             AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA,
             CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI,
             GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM,
             KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN,
             MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU,
             SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA,
             UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU,
             IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GO, GW, ML, MR, NE, SN, TD, TG,
             BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW,
             AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
     EP 1799205
                          Α1
                                20070627 EP 2005-752830
                                                                    200506
                                                                    13
             AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU,
             IE, IS, IT, LI, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR
     JP 2008506664
                          Τ
                                20080306
                                           JP 2007-520795
                                                                    200506
                                                                    13
     US 20080292562
                          Α1
                                20081127 US 2008-572199
                                                                    200807
                                                                    29
PRAI EP 2004-16878
                          Α
                                20040716
     WO 2005-EP52704
                          W
                                20050613
     The present invention relates to novel pharmaceutical compns. based
AΒ
     on PDE IV inhibitors and salts of glycopyrrolate salts, processes for
     preparing them and their use in the treatment of respiratory
     complaints. Thus, a formulation contained a glycopyrrolate salt 60,
     AWD 12281 200, lactose 12240 µg/capsule.
     873295-30-8
ΙT
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (pharmaceuticals for inhalation comprising PDE IV inhibitors and
        glycopyrrolate salts)
     873295-30-8 HCAPLUS
RN
     Pyrrolidinium, 3-[(2-cyclopentyl-2-hydroxy-2-phenylacetyl)oxy]-1,1-
CN
     dimethyl-, fluoride (1:1) (CA INDEX NAME)
```

• F -

```
CC
    63-6 (Pharmaceuticals)
ΙΤ
    56-81-5, Glycerol, biological studies 57-55-6, 1,2-Propanediol,
    biological studies 58-55-9, Theophylline, biological studies
    60-00-4, biological studies 64-02-8, Sodium Edetate
                                                         64-17-5,
    Ethanol, biological studies 67-63-0, 2-Propanol, biological
              596-51-0
                        25322-68-3 25322-68-3D, Polyethylene glycol,
                                       41078-02-8
                                                    135637-46-6
    esters with fatty acids 25322-69-4
                             146426-61-1 153259-65-5, Cilomilast
    136145-07-8, Arofylline
                 154284-39-6 155043-84-8
                                           161918-68-9 162278-09-3
    153587-17-8
    162401-32-3, Roflumilast 162542-90-7 179024-48-7, PD 168787
                 185954-27-2 185954-42-1
                                            186461-26-7, T 2585
    182282-60-6
    190377-71-0, NCS 613 192819-27-5 201932-04-9D, salts
                         207993-12-2, BY 343
    202185-74-8D, salts
                                              257892-34-5
    292135-78-5 329306-27-6 444659-35-2, YM 58997 444659-43-2
    475468-09-8
                 478409-36-8, CP 325366
                                         586388-32-1
                                                    746600-85-1D,
            754152-54-0D, salts 835882-98-9
                                              835882-99-0
    836652-75-6, C 1-1018 836652-82-5, CDC 3052
                                                  836652-83-6, Z 15370
    873295-30-8 873295-31-9 873295-32-0
                                            873295-33-1
    873295-34-2 873295-35-3
                               873295-36-4
                                            873295-37-5 873295-38-6
```

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (pharmaceuticals for inhalation comprising PDE IV inhibitors and glycopyrrolate salts)

873295-39-7 873295-40-0 873295-41-1 873295-43-3 873295-44-4

OSC.G 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L11 ANSWER 2 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN
- AN 2005:697566 HCAPLUS Full-text

873295-45-5 873295-46-6

- DN 144:78349
- TI Crystal structure of (3S,1'S)-2,2-dimethyl-3-[1,2-cyclohexylidenedioxyethyl]tetrahydro-1,2-oxazolium tetrafluoroborate, (C13H24NO3)[BF4]
- AU Frey, W.; Henneboehle, M.; Jaeger, V.

CS Institut fuer Organische Chemie, Universitaet Stuttgart, Stuttgart, 70569, Germany

SO Zeitschrift fuer Kristallographie - New Crystal Structures (2005), 220(2), 149-150

CODEN: ZKNSFT; ISSN: 1433-7266

PB Oldenbourg Wissenschaftsverlag GmbH

DT Journal

LA English

The title compound is orthorhombic, space group C2221, with a 9.320(2), b 10.6761(8), c 33.786(2) Å; Z = 8. Atomic coordinates are given. The isoxazolidine ring system shows an envelope conformation. Three atoms (C10, C11, C12) of the cyclohexylidene moiety exhibit very large displacement parameters.

(crystal structure of)

RN 773091-89-7 HCAPLUS

CN Isoxazolidinium, 3-(2S)-1,4-dioxaspiro[4.5]dec-2-yl-2,2-dimethyl-, (3S)-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 773091-88-6 CMF C13 H24 N O3

Absolute stereochemistry.

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

- CC 75-8 (Crystallography and Liquid Crystals) Section cross-reference(s): 28
- OSC.G 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)
- RE.CNT 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L11 ANSWER 3 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN
- AN 2005:356821 HCAPLUS Full-text
- DN 144:6402
- TI Kinetic study of the 5-exo cyclization of α -ammonium distonic radical cation, α -[N-(3-methyl-3-butenyl), N,N-(dimethyl)] ammoniomethyl
- AU Luz, Amalia Rios V.; Luz, Marina Jaramillo G.; Jaime, Martin F.
- CS Departamento de Quimica, Facultad de Ciencias Exactas y Naturales, Universidad de Caldas, Manizales, Colombia
- SO Revista Colombiana de Quimica (2004), 33(1), 21-31 CODEN: RCLQAY; ISSN: 0120-2804
- PB Universidad Nacional de Colombia, Departamento de Quimica
- DT Journal
- LA Spanish
- OS CASREACT 144:6402
- AB Cyclization of the α -ammonio distonic radical cation analogous to a 5-hexenyl system with Me substituent at 5 position and generated from α -[N-(3-methyl-3-butenyl), N,N-(dimethyl)] ammoniomethyl iodide via radical conditions, was studied by the unimol. vs. bimol. competition experiment Thus, under pseudo first order conditions, using nBu3SnH as reducing agent and photolytic initiation, the cyclization constant was determined as kc = 2.9 + 107, which means that the rate is very fast and the cyclization highly regionelective for the formation of pyrrolidinic ring systems substituted at 3 position, with recognized synthetic potential.
- IT 183849-69-6
 - RL: FMU (Formation, unclassified); PRP (Properties); FORM (Formation, nonpreparative)

(kinetic study of 5-exo cyclization of α -ammonium distonic radical cation, α -[N-(3-methyl-3-butenyl), N,N-(dimethyl)] ammoniomethyl)

RN 183849-69-6 HCAPLUS

CN Methyl, (1,1-dimethylpyrrolidinium-3-yl)phenyl-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 183849-68-5 CMF C13 H19 N

CM 2

CRN 14874-70-5

CMF B F4

CC 22-8 (Physical Organic Chemistry)

Section cross-reference(s): 74

IT 183849-69-6

RL: FMU (Formation, unclassified); PRP (Properties); FORM (Formation, nonpreparative)

(kinetic study of 5-exo cyclization of α -ammonium distonic radical cation, α -[N-(3-methyl-3-butenyl), N,N-(dimethyl)] ammoniomethyl)

OSC.G 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1

CITINGS)

RE.CNT 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 4 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 2004:392835 HCAPLUS Full-text

DN 141:332422

TI Synthesis of isoxazolines. 25. Isoxazolinium salts in asymmetric synthesis. 1. Stereoselective reduction induced by a 3'-alkoxy stereocenter. A new approach to polyfunctionalized β -amino acids

AU Henneboehle, Macro; Le Roy, Pierre-Yves; Hein, Matthias; Ehrler, Rudolf; Jaeger, Volker

CS Institut fuer Organische Chemie, Universitaet Stuttgart, Stuttgart, D-70569, Germany

SO Zeitschrift fuer Naturforschung, B: Chemical Sciences (2004), 59(4), 451-467

CODEN: ZNBSEN; ISSN: 0932-0776

PB Verlag der Zeitschrift fuer Naturforschung

DT Journal

LA English

OS CASREACT 141:332422

GΙ

AB A new approach to optically active N-methylamino acids was presented, relying on stereoselective reduction of N-methylisoxazolinium salts with a dioxyethyl side-chain. The diastereoselectivity of the reduction step was studied systematically, in comparison with that of resp. isoxazolines. A two-step transformation of isoxazolinium salts — with NaBH3(OAc) and subsequent catalytic hydrogenation as well as a one-pot reduction by catalytic hydrogenation led to high (95:5 and 87:13) diastereomeric ratios of protected erythro-N-methylaminopentanetriols. The hydroxyethyl side-chain was elaborated by oxidation to afford the β -N-methylamino acid I, exemplifying the potential of this strategy.

IT 773091-89-7P

RL: SPN (Synthetic preparation); PREP (Preparation) (asym. synthesis of N-methyl- β -amino acids and aminotriols via stereoselective reduction and stereoselective catalytic hydrogenation of N-methylisoxazolinium salts)

RN 773091-89-7 HCAPLUS

CN Isoxazolidinium, 3-(2S)-1, 4-dioxaspiro[4.5]dec-2-yl-2, 2-dimethyl-, (3S)-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 773091-88-6 CMF C13 H24 N O3

Absolute stereochemistry.

CM 2

CRN 14874-70-5

CMF B F4

CC 34-2 (Amino Acids, Peptides, and Proteins)

Section cross-reference(s): 28, 33

ΙT 110045-70-0P 771479-13-1P 771479-14-2P 771479-15-3P 771479-16-4P 771479-19-7P 771479-21-1P 771479-22-2P 771479-23-3P 771479-24-4P 771479-29-9P 771479-35-7P 773091-83-1P 773091-85-3P 773091-89-79

RL: SPN (Synthetic preparation); PREP (Preparation) (asym. synthesis of N-methyl- β -amino acids and aminotriols via stereoselective reduction and stereoselective catalytic hydrogenation of N-methylisoxazolinium salts)

RE.CNT 128 THERE ARE 128 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 5 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 2003:404002 HCAPLUS Full-text

DN 139:116999

TI New Insights in the Mechanism of Amine Catalyzed Epoxidation: Dual Role of Protonated Ammonium Salts as Both Phase Transfer Catalysts and Activators of Oxone

AU Aggarwal, Varinder K.; Lopin, Chrystel; Sandrinelli, Franck

CS School of Chemistry, University of Bristol, Bristol, BS8 1TS, UK

Journal of the American Chemical Society (2003), 125(25), 7596-7601 CODEN: JACSAT; ISSN: 0002-7863

PB American Chemical Society

DT Journal

LA English

OS CASREACT 139:116999

GΙ

Amines were previously reported to catalyze the epoxidn. of alkenes using Oxone (2KHSO5+KHSO4+K2SO4), and significant levels of asym. induction were observed From screening amines based on 2-substituted pyrrolidines, it has now been found that more consistent and reproducible results are achieved with the HCl salt of the amine compared to the amine itself. Up to 66% ee was achieved in epoxidn. of 1-phenylcyclohexene. The chiral amine could be reisolated in >90% yield when reactions were conducted at -10°, indicating that the integrity of the amine was maintained during the oxidation process. At -10°, (S)-2-(diphenylmethyl)pyrrolidine 1 reacted with Oxone to give a mixture of ammonium salts containing the peroxymonosulfate salt I. The enantioselectivity obtained with this salt was compared to the amine·HCl salt catalyzed process and identical results were observed, indicating that the true oxidant was the peroxymonosulfate

salt I. The relative rates of oxidation of cis- and trans- β -methylstyrenes together with the ρ value of 1-arylcyclohexenes were determined. The amine catalyzed process involved electrophilic oxidation. A new mechanism is advanced in which the protonated amine not only acts as a PTC but also activates Oxone, through H bonding, toward electrophilic attack.

IT 562813-65-4P

RL: CAT (Catalyst use); CPS (Chemical process); PEP (Physical, engineering or chemical process); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); PROC (Process); RACT (Reactant or reagent); USES (Uses)

(nonstereoselective low yield 1-phenylcyclohexene epoxidn. by; dual role of protonated ammonium salts as both phase transfer catalysts and activators of oxone in mechanism of amine catalyzed epoxidn.)

RN 562813-65-4 HCAPLUS

CN Pyrrolidinium, 2-(diphenylmethyl)-1,1-dimethyl-, (2S)-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 562813-64-3 CMF C19 H24 N

Absolute stereochemistry. Rotation (+).

CM 2

CRN 14874-70-5

CMF B F4

$$-F = \begin{bmatrix} F & -1 \\ 3 & 3 + \end{bmatrix}$$

```
CC 22-3 (Physical Organic Chemistry)
```

IT 562813-65-4P

RL: CAT (Catalyst use); CPS (Chemical process); PEP (Physical, engineering or chemical process); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); PROC (Process); RACT (Reactant or reagent); USES (Uses)

(nonstereoselective low yield 1-phenylcyclohexene epoxidn. by; dual role of protonated ammonium salts as both phase transfer catalysts and activators of oxone in mechanism of amine catalyzed epoxidn.)

- OSC.G 40 THERE ARE 40 CAPLUS RECORDS THAT CITE THIS RECORD (42 CITINGS)
- RE.CNT 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L11 ANSWER 6 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN
- AN 2002:889558 HCAPLUS Full-text
- DN 137:369966
- TI Preparation of enantiomerically pure basic [(cyclopentyl- or cyclohexylhydroxyphenylacetyl)oxy]-1,1-dimethylpyrrolidinium salts, their muscarinic receptor binding affinity, and use as treatment for obstructive respiratory disease
- IN Noe, Christian; Mutschler, Ernst; Lambrecht, Gunter; Elgert, Michael; Elgert, Ruth Irene; Czeche, Sittah; Waelbroeck, Magali
- PA Germany
- SO U.S. Pat. Appl. Publ., 19 pp., Cont.-in-part of U.S. 6,307,060. CODEN: USXXCO
- DT Patent
- LA English
- FAN.CNT 2

r AN.	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE -
ΡΙ	 US 20020173536	A1	20021121	US 2001-901217	200107
	US 6613795 WO 9821183	B2 A1	20030902 19980522	WO 1997-AT245	09
		300 311 36			199711 11

W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR,

TT, UA, UG, US, UZ, VN, YU, ZW RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG EP 2003-5233 20031210 EP 1369414 Α1 199711 11 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, AL 20031217 EP 1371645 Α1 EP 2003-5232 199711 11 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, AL 20011023 US 6307060 В1 US 1999-309960 199905 11 US 20030220400 A1 20031127 US 2003-601542 200306 23 US 7253182 В2 20070807 PRAI AT 1996-1973 Α 19961111 WO 1997-AT245 W 19971111 US 1999-309960 Α2 19990511 EP 1997-911049 Α3 19971111 US 2001-901217 20010709 ΑЗ ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OS MARPAT 137:369966

Ι

GΙ

AB Disclosed are enantiomerically pure cyclic aminoalc. esters of arylcycloalkylhydroxycarboxylic acids with at least 90% enantiomeric excess of the (3R,2'R), (3S,2'R), (3R,2'S), or (3S,2'S) configured

enantiomer. Thus, [(cyclopentyl (or cyclohexyl) hydroxyphenylacetyl)oxy] pyrrolidinium salts I (R = cyclopentyl, cyclohexyl, X = bromide, iodide, fluoride, chloride) were prepared by reacting (3R) or (3S)-1-methyl-3-pyrrolidinol with the corresponding phenylacetate, followed by preparation of the tartrate intermediates and quaternization. Inhalable powder and aerosol formulations of the compds. were also prepared The muscarinic binding affinity of I were examined using rabbit vas deferens, guinea pic atrium, guinea pig ileum, and human M1, M2, and M3 receptors.

IT 475468-10-1P 475468-11-2P 475468-17-8P 475468-23-6P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

 $\hbox{ (preparation and muscarinic receptor binding affinity of } \\ [\hbox{ (cyclopentyl } \\$

or cyclohexylhydroxyphenylacetyl)oxy]pyrrolidinium halides and inhalable and aerosol formulation prepns. for treating obstructive respiratory diseases)

RN 475468-10-1 HCAPLUS

CN Pyrrolidinium, 3-[[(2R)-2-cyclopentyl-2-hydroxy-2-phenylacetyl]oxy]-1,1-dimethyl-, fluoride (1:1), (3R)- (CA INDEX NAME)

Absolute stereochemistry.

● F -

RN 475468-11-2 HCAPLUS

CN Pyrrolidinium, 3-[[(2R)-2-cyclopentyl-2-hydroxy-2-phenylacetyl]oxy]-1,1-dimethyl-, fluoride (1:1), (3S)- (CA INDEX NAME)

Absolute stereochemistry.

RN 475468-17-8 HCAPLUS

CN Pyrrolidinium, 3-[[(2R)-2-cyclohexyl-2-hydroxy-2-phenylacetyl]oxy]-1,1-dimethyl-, fluoride (1:1), (3S)- (CA INDEX NAME)

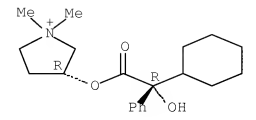
Absolute stereochemistry.

• F-

RN 475468-23-6 HCAPLUS

CN Pyrrolidinium, 3-[[(2R)-2-cyclohexyl-2-hydroxy-2-phenylacetyl]oxy]-1,1-dimethyl-, fluoride (1:1), (3R)- (CA INDEX NAME)

Absolute stereochemistry.



• F-

```
IC
    ICM A61K031-4015
    ICS C07D207-46
INCL 514424000; X54-854.2
    27-10 (Heterocyclic Compounds (One Hetero Atom))
CC
    Section cross-reference(s): 63
    129784-11-8P
                   129784-12-9P
                                 129784-14-1P
                                                 201667-20-1P
ΙT
                   207856-75-5P 207856-76-6P
    207856-74-4P
                                                 207856-77-7P
    207856-78-8P
                   207856-79-9P
                                                 207856-81-3P
                                  207856-80-2P
    475468-09-8P 475468-10-1P 475468-11-2P
    475468-13-4P 475468-15-6P
                                  475468-17-8P
                                                 475468-19-0P
                                  475468-25-8P
    475468-21-4P
                   475468-23-6P
    RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
```

or cyclohexylhydroxyphenylacetyl)oxy]pyrrolidinium halides and inhalable and aerosol formulation prepns. for treating obstructive respiratory diseases)

OSC.G 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS)

L11 ANSWER 7 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 2001:174677 HCAPLUS Full-text

DN 134:366563

- TI Construction of Persistent Phenoxyl Radical with Intramolecular Hydrogen Bonding
- AU Maki, Toshihide; Araki, Yoko; Ishida, Yukihiro; Onomura, Osamu; Matsumura, Yoshihiro
- CS Faculty of Pharmaceutical Sciences, Nagasaki University, Nagasaki, 852-8521, Japan
- Journal of the American Chemical Society (2001), 123(14), 3371-3372 CODEN: JACSAT; ISSN: 0002-7863

PB American Chemical Society

DT Journal

LA English

OS CASREACT 134:366563

GI

AB The authors characterized phenoxy radical cation I as a proper model for H-bonded phenoxy radicals in biol. systems. In this model, an intramol. proton migration occurs at its redox process with the min. nuclear motion requirement. Probably some basic functional group located near the tyrosine residue in an active site of enzyme may control the redox potential of the tyrosyl radical with H bonding.

IT 340228-55-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthetic precursor of 2-aminomethyl-4,6-tert-butylphenol analog; construction of persistent phenoxyl radical with intramol. hydrogen bonding)

RN 340228-55-9 HCAPLUS

CN Pyrrolidinium, 2-[3,5-bis(1,1-dimethylethyl)-2-hydroxyphenyl]-1,1-dimethyl-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 340228-54-8 CMF C20 H34 N O

CM 2

CRN 14874-70-5

CMF B F4

CCI CCS

CC 22-12 (Physical Organic Chemistry)

Section cross-reference(s): 7, 34, 72

IT 340228-55-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthetic precursor of 2-aminomethyl-4,6-tert-butylphenol analog; construction of persistent phenoxyl radical with

intramol. hydrogen bonding)

OSC.G 44 THERE ARE 44 CAPLUS RECORDS THAT CITE THIS RECORD (47 CITINGS)

RE.CNT 34 THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 8 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1999:532438 HCAPLUS Full-text

DN 131:299336

TI Triarylaminium Salt Induced Oxidative Cyclizations of Tertiary Amines. Convenient Access to 2-Substituted Pyrrolidinium Salts

AU Jahn, Ullrich; Aussieker, Susanne

CS Institut fuer Organische Chemie, Technische Universitaet Braunschweig, Braunschweig, D-38106, Germany

SO Organic Letters (1999), 1(6), 849-852 CODEN: ORLEF7; ISSN: 1523-7060

PB American Chemical Society

DT Journal

LA English

OS CASREACT 131:299336

AB Convenient oxidative generation and facile 5-exo cyclization of tertiary aminium radical cations to give distonic 2-substituted pyrrolidinium radical cations is reported. These can be further oxidized to 1,3-dications and trapped by nucleophiles as water, alcs., or chloride ion. Preliminary mechanistic issues and implications will be presented.

IT 247118-21-4P 247118-27-0P

247118-31-6P 247118-35-0P 247118-37-2P 247118-39-4P 247118-40-7P 247118-43-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (triarylaminium salt induced oxidative cyclization of tertiary amines)

RN 247118-21-4 HCAPLUS

CN Pyrrolidinium, 1,1-diethyl-2-(hydroxydiphenylmethyl)-, hexafluorophosphate(1-) (9CI) (CA INDEX NAME)

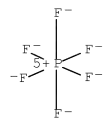
CM 1

CRN 247118-20-3 CMF C21 H28 N O

CM 2

CRN 16919-18-9

CMF F6 P CCI CCS

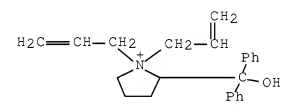


RN 247118-27-0 HCAPLUS

CN Pyrrolidinium, 2-(hydroxydiphenylmethyl)-1,1-di-2-propen-1-yl-, hexafluorophosphate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 247118-26-9 CMF C23 H28 N O



CM 2

CRN 16919-18-9

CMF F6 P CCI CCS

RN 247118-31-6 HCAPLUS

CN Pyrrolidinium, 1,1-diethyl-2-(hydroxyphenylmethyl)-, hexafluorophosphate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 247118-30-5 CMF C15 H24 N O

CM 2

CRN 16919-18-9

CMF F6 P CCI CCS

RN 247118-35-0 HCAPLUS

CN Pyrrolidinium, 1,1-diethyl-2-(methoxydiphenylmethyl)-, hexafluorophosphate(1-) (9CI) (CA INDEX NAME)

CM 1

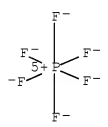
CRN 247118-34-9

CMF C22 H30 N O

CM 2

CRN 16919-18-9

CMF F6 P CCI CCS



RN 247118-37-2 HCAPLUS

CN Pyrrolidinium, 1,1-diethyl-2-(methoxyphenylmethyl)-, hexafluorophosphate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 247118-36-1 CMF C16 H26 N O

CM 2

CRN 16919-18-9

CMF F6 P CCI CCS

RN 247118-39-4 HCAPLUS

CN Pyrrolidinium, 2-(ethoxydiphenylmethyl)-1,1-diethyl-, hexafluorophosphate(1-) (9CI) (CA INDEX NAME)

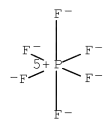
CM 1

CRN 247118-38-3 CMF C23 H32 N O

CM 2

CRN 16919-18-9

CMF F6 P CCI CCS

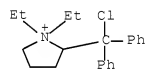


RN 247118-40-7 HCAPLUS

CN Pyrrolidinium, 2-(chlorodiphenylmethyl)-1,1-diethyl-, hexafluorophosphate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 247118-22-5 CMF C21 H27 C1 N



CM 2

CRN 16919-18-9

CMF F6 P CCI CCS

RN 247118-43-0 HCAPLUS
CN Pyrrolidinium, 2-benzoyl-1,1-diethyl-, hexafluorophosphate(1-) (9CI)
(CA INDEX NAME)

CM 1

CRN 247118-42-9
CMF C15 H22 N O

CM 2

CRN 16919-18-9

CMF F6 P

CCI CCS

27-10 (Heterocyclic Compounds (One Hetero Atom)) CC 247118-21-4P ΙT 247118-23-6P 247118-25-8P 247118-27-0P 247118-29-2P **247118-31-6P** 247118-33-8P 247118-35-0P 247118-37-2P 247118-39-4P 247118-40-7P 247118-41-8P 247118-43-0P 247118-45-2P 308140-71-8P 308141-00-6P RL: SPN (Synthetic preparation); PREP (Preparation) (triarylaminium salt induced oxidative cyclization of tertiary amines)

- OSC.G 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)
- RE.CNT 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L11 ANSWER 9 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN
- AN 1996:637151 HCAPLUS Full-text
- DN 125:339949
- OREF 125:63349a,63352a
- TI Generation and Study of the Reactivity of $\alpha\text{--Ammonium Distonic}$ Radical Cations in Solution
- AU Rios, Luz Amalia; Dolbier, William R. Jr.; Paredes, Rodrigo; Lusztyk, Janusz; Ingold, K. U.; Jonnson, Mats
- CS Department of Chemistry, University of Florida, Gainesville, FL, 32611-7200, USA
- SO Journal of the American Chemical Society (1996), 118(45), 11313-11314

 CODEN: JACSAT; ISSN: 0002-7863
- PB American Chemical Society
- DT Journal
- LA English
- AB α -Ammonium distonic radical cations have for the first time been purposefully generated and their reactivity studied in solution using both LFP and competition expts. Such species exhibit behavior which is typical of a carbon-bound radical, and rate consts. of cyclization of unsubstituted-, 5-phenyl-, 5,5-diphenyl-4-pentyldimethylammoniomethyl radicals (1.5 + 106 s-1 at 33°, 1.7 + 107 s-1 at 25°, and 3.0 + 107 s-1 at 25° at 33°) indicated that they are somewhat more reactive than analogous hydrocarbon radicals.
- IT 183849-69-6
 - RL: FMU (Formation, unclassified); PEP (Physical, engineering or chemical process); PRP (Properties); FORM (Formation, nonpreparative); PROC (Process)

(generation and study of reactivity of $\alpha\text{-ammonium}$ distonic radical cations in solution)

- RN 183849-69-6 HCAPLUS
- CN Methyl, (1,1-dimethylpyrrolidinium-3-yl)phenyl-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 183849-68-5 CMF C13 H19 N

CM 2

CRN 14874-70-5

CMF B F4

CCI CCS

CC 67-3 (Catalysis, Reaction Kinetics, and Inorganic Reaction Mechanisms)

Section cross-reference(s): 22

IT 661-36-9 183849-67-4 **183849-69-6** 183849-71-0 183849-73-2 183849-75-4 183849-77-6 183849-79-8 183849-81-2 183849-83-4

RL: FMU (Formation, unclassified); PEP (Physical, engineering or chemical process); PRP (Properties); FORM (Formation, nonpreparative); PROC (Process)

(generation and study of reactivity of α -ammonium distonic radical cations in solution)

OSC.G 14 THERE ARE 14 CAPLUS RECORDS THAT CITE THIS RECORD (14 CITINGS)

L11 ANSWER 10 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1995:792939 HCAPLUS Full-text

DN 123:354687

OREF 123:63367a,63370a

TI Heat sensitive color developing material

IN Tanabe, Hisake; Nakano, Shingi; Nakae, Yasuhiko; Urano, Satoshi; Eguchi, Yoshio

PA Nippon Paint Co., Ltd., Japan

SO U.S., 11 pp. Cont.-in-part of U.S. Ser. No. 727,671, abandoned.

CODEN: USXXAM

DT Patent

LA English

FAN.CNT 2

T 1214 •	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE –
ΡI	 US 5439516	A	19950808	US 1993-175692	199312
	JP 04067988	A	19920303	JP 1990-181878	30
PRAI	: JP 1990-181878	A	19900709		09

PRAI JP 1990-181878 US 1991-727671

B2 19900709

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OS MARPAT 123:354687

AB A heat sensitive color developing material containing an electron donating color forming organic compound, a heat activating compound, and a color developer. This material exhibits excellent heat sensitivity and enables rapid color development at low cost.

IT 144382-76-3 171101-69-2

RL: TEM (Technical or engineered material use); USES (Uses) (heat sensitive color developing material)

RN 144382-76-3 HCAPLUS

CN Oxazolidinium, 3,3-dimethyl-2-phenyl-, (OC-6-11)-, hexafluoroantimonate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 144382-75-2 CMF C11 H16 N O

CM 2

CRN 17111-95-4

171101-69-2 HCAPLUS RN

Oxazolidinium, 3-ethyl-2-(4-methoxyphenyl)-3-methyl-, CN hexafluorophosphate(1-) (1:1) (CA INDEX NAME)

CM 1

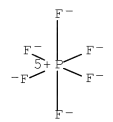
CRN 171101-68-1 CMF C13 H20 N O2

CM2

CRN 16919-18-9

CMF F6 P

CCI CCS



GΙ

```
IC
     ICM C09D011-00
INCL 106-21R
CC
     74-6 (Radiation Chemistry, Photochemistry, and Photographic and
     Other Reprographic Processes)
     541-16-2, Di-tert-butyl malonate 20270-53-5, Di-tert-butyl adipate
ΙT
     20487-40-5, tert-Butyl propionate 25852-37-3, Butyl
     acrylate-methyl methacrylate copolymer 29035-74-3, Butyl
     acrylate-butyl methacrylate copolymer 29535-38-4, Butyl
                                                  50292-91-6
     methacrylate-lauryl methacrylate copolymer
                                                               75805-17-3
     89331-94-2 133227-06-2 136608-77-0 144382-74-1
     144382-76-3
                  144382-80-9 144382-82-1 144382-85-4
     171101-69-2
     RL: TEM (Technical or engineered material use); USES (Uses)
        (heat sensitive color developing material)
              THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 11 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN
L11
ΑN
     1994:508979 HCAPLUS Full-text
     121:108979
DN
OREF 121:19691a,19694a
     Synthesis and Rearrangement of Intramolecularly Stabilized
ΤI
     1\sigma^2, 3\sigma^2-Diphosphaallylic Cations into Intramolecularly
     Stabilized 1\sigma1, 3\sigma3-Diphosphaallylic Cations
     Soleilhavoup, Michele; Canac, Yves; Polozov, Alexander M.;
ΑU
     Baceiredo, Antoine; Bertrand, Guy
CS
     Laboratoire de Chimie de Coordination, CNRS, Toulouse, 31077, Fr.
SO
     Journal of the American Chemical Society (1994), 116(14), 6149-52
     CODEN: JACSAT; ISSN: 0002-7863
DT
     Journal
    English
LA
     CASREACT 121:108979
OS
```

AΒ Two equivalent of boron trifluoride-diethyl ether complex or 1 equiv of trifluoromethanesulfonic acid reacts with C-[bis(diisopropylamino)phosphino]-C,Pbis(diisopropylamino)phosphaalkene R2NP:C(NR2)P(NR2)2 (R = Me2C), cleaving one of the diisopropylamino substituents at the σ 3phosphorus atom. This affords the corresponding intermediate $1\sigma^2$, $3\sigma^2$ -diphosphaallylic cation I, which is isolated as the fourmembered-heterocycle II. Addition of a catalytic amount of base to II leads to a $1\sigma 1$, $3\sigma 3$ -diphosphaallylic cation intermediate III, which is isolated as the diphosphirenium salt IV. Mesityllithium and the lithium salts of diisopropylamine or dicyclohexylamine react at the σ 2-phosphorus atom of the cationic heterocycles II and IV, affording the corresponding phosphaalkenes. All the results demonstrate the high electrophilicity of low-coordinated diphosphaallylic cations, which are only isolable as intramol. donor-acceptor complexes.

IT 156664-17-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction with base)

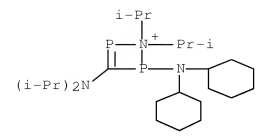
RN 156664-17-4 HCAPLUS

1,2,4-Azadiphosphetium, 3-[bis(1-methylethyl)amino]-2(dicyclohexylamino)-1,1-bis(1-methylethyl)-, tetrafluoroborate(1-)
(1:1) (CA INDEX NAME)

CM 1

CN

CRN 156664-16-3 CMF C25 H50 N3 P2



CRN 14874-70-5

CMF B F4

CC 29-7 (Organometallic and Organometalloidal Compounds)

Section cross-reference(s): 22

IT 156664-15-2P 156664-17-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction with base)

OSC.G 20 THERE ARE 20 CAPLUS RECORDS THAT CITE THIS RECORD (20 CITINGS)

L11 ANSWER 12 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1994:446709 HCAPLUS Full-text

DN 121:46709

OREF 121:8259a,8262a

TI Heat-sensitive color-developing recording material

IN Tanabe, Hisaki; Nakae, Yasuhiko; Nakano, Shingi; Eguchi, Yoshio

PA Nippon Paint Co., Ltd., Japan

SO U.S., 15 pp. Division of U.S. Ser. No. 726,835. CODEN: USXXAM

DT Patent

LA English

10/536,829

F	'AN	CNT	2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	 US 5288686	 А	19940222	US 1993-17481	
	US 53021 9 4	A	19940412	US 1991-726835	199302 12
	1001 50005	- 0			199107 08
PRAI	US 1991-726835 JP 1990-181877	A3 A	19910708 19900709		

AB A heat-sensitive color-developing recording material contains an electron-donating color-forming organic compound, a heat-activating compound, and a color developer. The material has excellent heat sensitivity in which color can be developed rapidly.

IT 144382-76-3

RL: USES (Uses)

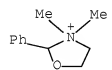
(heat-sensitive color-developing recording materials containing)

RN 144382-76-3 HCAPLUS

CN Oxazolidinium, 3,3-dimethyl-2-phenyl-, (OC-6-11)-, hexafluoroantimonate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 144382-75-2 CMF C11 H16 N O

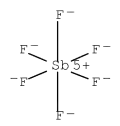


CM 2

CRN 17111-95-4

CMF F6 Sb

CCI CCS



```
IC ICM B41M005-20 ICS C09D011-02
```

INCL 503209000

CC 74-7 (Radiation Chemistry, Photochemistry, and Photographic and Other Reprographic Processes)

IT 20270-53-5, Di-tert-butyl adipate 31604-40-7 50292-91-6, 3,3-Bis(1-butyl-2-methylindol-3-yl)phthalide 64309-46-2 71436-70-9 75805-17-3 89331-94-2,

3-Dibutylamino-6-methyl-7-(phenylamino)fluoran 129254-44-0 131212-08-3 136608-77-0 144031-48-1 144031-49-2 144382-74-1

144382-76-3144382-80-9144382-84-3144382-85-4147341-81-9147360-96-1156083-92-0156083-93-1

RL: USES (Uses)

(heat-sensitive color-developing recording materials containing)

OSC.G 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 13 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1994:220511 HCAPLUS Full-text

DN 120:220511

OREF 120:39141a,39144a

TI Two-coat-one-bake process for solvent-based high-solids metallic coating

IN Nakano, Shinji; Nishizawa, Koji; Okude, Yoshitaka

PA Nippon Paint Co Ltd, Japan

SO Jpn. Kokai Tokkyo Koho, 19 pp. CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

	01.1				
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PΙ	JP 05253537	A	19931005	JP 1992-89907	

JP 2844280 PRAI JP 1992-89907 В2 19990106 19920312

The title process providing high-performance coatings involves AΒ application of metallic pigment-containing base coat and clear topcoat, containing curing catalysts containing onium salts comprising (non)cyclic ammonium cation or sulfonium cation with R1R2R3C6H2C(R4)(R5) group bonded to N or S and AsF-6, SbF-6, BF-4, PF-6, ClO-4, FeCl-4, CF3SO-3, aromatic or aliphatic sulfonic or carboxylic acid ion. Acrylic or polyester vehicles were used with acrylic microgels, aminoplast hardeners, Al pigments, and curing catalysts, e.g., 1-(4-methylbenzyl)tetrahydrothiophenium hexafluoroantimonate, N-(2,6-dichlorobenzyl)-4-cyanopyridinium ptoluenesulfonate, N-(4-chlorobenzyl)-N, N-dimethylanilinium 4dodecylbenzenesulfonate, N-(2-methylbenzyl)-4-cyanopyridinium hexafluorophosphate, etc., at coating solids content ≥45%.

154396-39-1 ΙT

> RL: CAT (Catalyst use); USES (Uses) (crosslinking catalysts, for acrylic and polyester coatings containing aminoplasts)

154396-39-1 HCAPLUS RN

Oxazolium, 2,3-dihydro-3,3-dimethyl-2-phenyl-, CN hexafluorophosphate(1-) (1:1) (CA INDEX NAME)

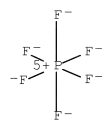
CM 1

CRN 141915-33-5 CMF C11 H14 N O

CM 2

16919-18-9 CRN

CMF F6 P CCI CCS



ΙC ICM B05D005-06

ICS B05D001-36; B05D005-06; B05D007-24

CC 42-10 (Coatings, Inks, and Related Products)

97744-46-2 153359-44-5 153359-45-6 153359-48-9 153359-49-0 ΙT

153359-50-3 153359-51-4 153359-53-6 154396-39-1

154396-40-4 154397-34-9

RL: CAT (Catalyst use); USES (Uses)

(crosslinking catalysts, for acrylic and polyester coatings containing aminoplasts)

L11 ANSWER 14 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1993:505987 HCAPLUS Full-text

DN 119:105987

OREF 119:18875a,18878a

Heat-sensitive coloring material for recording sheet TΙ

IN Tanabe, Hisanori; Nakae, Yasuhiko; Nakano, Shinji; Eguchi, Yoshio

PA Nippon Paint K. K., Japan

SO Jpn. Kokai Tokkyo Koho, 12 pp.

CODEN: JKXXAF

DT Patent

LAJapanese

FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	JP 04067987	A	19920303	JP 1990-181877	199007
	US 5302194	A	19940412	us 1991-726835	09 199107
PR A I	JP 1990-181877	A	19900709		08

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

GΙ

AΒ A heat-sensitive coloring material contains (A) an electron-donating coloring organic compound, (B) a heat-activating agent selected from cyclic sulfonium salts [I; R1 = H, R, OR, halo, NO2; R = (OHsubstituted) C1-12 alkyl, cycloalkyl; X = AsF6, SbF6, BF4, BF6, PF6, C104, FeC14, CF3S03, RS03, RC02], (R2)3C6H2C(R3)2A+X- [R2 = COR, R1, R2]OH, cyano, NH2; R3 = H, R, halo; A = Q, N(R4)3; when A = N(R4)3, at least one of R3 \neq H; R4 = (un)substituted C1-12 alkyl, alkenyl, or Ph; R, R1, X are defined as above], and heterocyclic ammonium salts [II; R5 = H, R, C2-3 alkenyl, R8; R6 = R, C2-3 alkenyl, R8; R7 = H, OH, R, OR, R8; R8 = (un) substituted Ph; m = 1-4; R, X are defined as above], and (C) a color-developing agent selected from aromatic compds.(III; Ar = benzene or naphthalene ring; R9 = H, OR, O2CR, COR, halo, NO2; R is defined as above) and diphenylmethane derivs. (IV; R is defined as above), and (D) a resin having number-average mol. weight 500-50,000. The recording material is not colored in the presence of an organic solvent, thus is easy to handle, and also the choice of heat-activating agent (B) allows the recording material to be rapidly colored at desired, relatively low temperature Furthermore only a very small amount of the expensive heat-activating agent is required to effectively initiate the decomposition of the tert-butoxyphenyl group to hydroxyphenyl group in the coloring agent and subsequent coloration, and thus the recording material is economically manufactured

IT 144382-76-3

RL: USES (Uses)

(heat-activating agent, heat-sensitive recording material

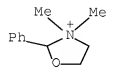
containing)

RN 144382-76-3 HCAPLUS

CN Oxazolidinium, 3,3-dimethyl-2-phenyl-, (OC-6-11)-, hexafluoroantimonate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 144382-75-2 CMF C11 H16 N O



CM 2

CRN 17111-95-4

CMF F6 Sb

IC ICM B41M005-26

ICS B41M005-30

CC 74-12 (Radiation Chemistry, Photochemistry, and Photographic and Other Reprographic Processes)

IT 126888-77-5 136608-77-0 144382-71-8 144382-74-1 144382-85-4 144382-85-4

RL: USES (Uses)

(heat-activating agent, heat-sensitive recording material containing)

OSC.G 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)

L11 ANSWER 15 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1993:437599 HCAPLUS Full-text

DN 119:37599

OREF 119:6695a,6698a

TI Heat-sensitive coloring material for recording sheet

IN Tanabe, Hisanori; Nakano, Shinji; Nakae, Yasuhiko; Urano, Satoru; Eguchi, Yoshio

PA Nippon Paint K. K., Japan

SO Jpn. Kokai Tokkyo Koho, 12 pp.

CODEN: JKXXAF

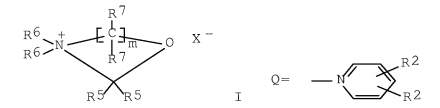
DT Patent

LA Japanese

FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	JP 04067988	A	19920303	JP 1990-181878	
					199007
					09
	US 5439516	A	19950808	US 1993-175692	
					199312
					30
PRAI	JP 1990-181878	A	19900709		
	US 1991-727671	B2	19910709		
7 C C T /	CNIMENIE HITOTODA DOD II		T 7777 T 7 D T D	TNI TOUR DECDIAN HODMAH	

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT GI



AB A heat-sensitive recording material contains (A) an electron-donating coloring organic compound, (B) a heat-activating agent selected from (R2)3C6H2C(R3)2A+X- [R2 = COR, R1, OH, cyano, NH2; R3 = H, R, halo; A

= Q, N(R4)3; when A = N(R4)3, at least one of $R3 \neq H$; R4 =(un) substituted C1-12 alkyl, alkenyl, or Ph; R = C1-4(hydroxy)(cyclo)alkyl; R1 = H, R, OR, halo, NO2; X = AsF6, SbF6, BF4, BF6, PF6, ClO4, FeCl4, CF3SO3, RSO3, RCO2] and heterocyclic ammonium salts [I; R5 = H, R, C2-3 alkenyl, R8; R6 = R, C2-3 alkenyl, R8; R7 =H, OH, R, OR, R8; R8 = (un)substituted Ph; m = 1-4; R, X are defined as above]. It addnl. contains a color developer, having higher b.p. than that of the heat-activating agent, selected from tert-Bu esters of C3-18 aliphatic carboxylic acids, C6-10 carbocyclic or heterocyclic carboxylic acids, and film-forming polymers having tert-Bu ester side chain and number-average mol. weight 500-50,000. recording material is not colored in the presence of an organic solvent, thus is easy to handle, and also the choice of heatactivating agent (B) allows the recording material to be rapidly colored at desired, relatively low temperature Furthermore only a very small amount of the expensive heat-activating agent is required to effectively initiate the decomposition of the tert-butoxyphenyl group to hydroxyphenyl group in the coloring agent and subsequent coloration, and thus the recording material is economically manufactured

IT 144382-76-3

RL: USES (Uses)

(heat-activating agent, heat-sensitive recording material containing)

RN 144382-76-3 HCAPLUS

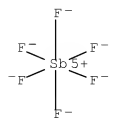
CN Oxazolidinium, 3,3-dimethyl-2-phenyl-, (OC-6-11)-, hexafluoroantimonate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 144382-75-2 CMF C11 H16 N O

CM 2

CRN 17111-95-4 CMF F6 Sb CCI CCS



IC ICM B41M005-26

CC 74-12 (Radiation Chemistry, Photochemistry, and Photographic and Other Reprographic Processes)

IT 133227-06-2 144382-70-7 144382-71-8 144382-72-9 144382-73-0

144382-74-1 144382-76-3 144382-78-5 144382-80-9

144382-82-1

RL: USES (Uses)

(heat-activating agent, heat-sensitive recording material containing)

L11 ANSWER 16 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1993:61658 HCAPLUS Full-text

DN 118:61658

OREF 118:11017a,11020a

TI Acrylic epoxy thermosetting resin compositions

IN Yamada, Atsushi; Takae, Masaki; Mure, Shoichi; Takeoka, Kazuhiko; Nakano, Shinji

PA Nippon Paint Co., Ltd., Japan

SO Jpn. Kokai Tokkyo Koho, 9 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	 JP 04139280	A	19920513	JP 1990-261978	
					199009 28

PRAI JP 1990-261978 19900928

AB Title compns. with good storage stability comprise (A) polymers having epoxy value 70-250, OH value <150, acid value <20, and number-average mol. weight 1000-20,000, (B) polyepoxy compds. and (C) heat-

sensitive latent cationic polymerization catalysts at A/B ratio 100:0-150 and C/(A + B) 0.05-5%. Thus, a clean composition cong. 0.5 part N-(4-chlorobenzyl)-N,N-dimethylanilinium SbF6- and 153.8 parts styrene-glycidyl methacrylate-2-hydroxyethyl methacrylate-Bu acrylate-Bu methacrylate copolymer showed good storage stability (50°, 10 days) and was baked at 140° for 20 min to give a film with good abrasion, chemical, water, and weather resistance.

IT 144382-76-3

RL: CAT (Catalyst use); USES (Uses)
(latent catalyst, heat-sensitive, for cationic polymerization, for acrylic epoxy coatings)

RN 144382-76-3 HCAPLUS

CN Oxazolidinium, 3,3-dimethyl-2-phenyl-, (OC-6-11)-, hexafluoroantimonate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 144382-75-2 CMF C11 H16 N O

CM 2

CRN 17111-95-4 CMF F6 Sb CCI CCS

$$F - \begin{cases} F - \\ Sb5 + F - \\ F - \end{cases}$$

- IC ICM C09D163-00 ICS B05D001-36; B05D007-24; C08G059-68; C08L063-00; C09D163-00 CC 42-9 (Coatings, Inks, and Related Products) 126803-31-4 136608-77-0 144207-48-7 144382-76-3 ΙΤ RL: CAT (Catalyst use); USES (Uses) (latent catalyst, heat-sensitive, for cationic polymerization, for acrylic epoxy coatings) L11 ANSWER 17 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN 1992:591985 HCAPLUS ΑN Full-text DN 117:191985 OREF 117:33171a,33174a Reaction of aminocarbene complexes of chromium with alkynes. 1. ΤI Formation and rearrangement of ketene and nitrogen ylide complexes Chelain, Evelyne; Goumont, Regis; Hamon, Louis; Parlier, Andree; ΑU Rudler, Michele; Rudler, Henri; Daran, Jean Claude; Vaissermann, Jacqueline CS Lab. Chim. Org., Univ. Pierre et Marie Curie, Paris, 75252, Fr. Journal of the American Chemical Society (1992), 114(21), 8088-98 SO CODEN: JACSAT; ISSN: 0002-7863 DT Journal LAEnglish CASREACT 117:191985 OS AB The title reactions of chromium-containing carbene complexes (CO) 5Cr:C(R1)N(R2R3) [R1 = H, Me, Ph; R2 = Me; R3 = Me, cyclopropyl, cyclopropylmethyl; R2R3 = (CH2)5] 8 and
- (CO) 5Cr:C[(CH2) 3C.tplbond.CPh]N(R1R2) [R1 = R2 = Me; R1R2 = (CH2)5, (CH2)4] 9, bearing alkyl groups of low migratory aptitude on nitrogen were examined In contrast to complexes in which nitrogen bears either an alkyl and an allyl or a benzyl group or is part of a strained cycle, which give heterocycles upon alkyne/CO insertions followed by nitrogen-to-carbon migrations, complexes 8 and 9 lead to stable nitrogen ylides, which could be fully characterized by x-ray crystallog. in the case of 8 [R1 = H, R2R3 = (CH2)5] and 9 (R1 = R2 =Me). Moreover, in the case of complexes of the general structure 9, ketene precursors of the ylides could either be detected (R1 = Me, R2 = CH2Ph) or isolated and characterized [R2R3 = (CH2)5]. ylide complexes undergo, upon moderate heating, Stevens-type rearrangements to the expected heterocyclic compds. as a result of nitrogen-to-carbon migrations of various alkyl groups, and upon treatment with dimethyldioxirane, they undergo oxidation to lactone complexes.
- IT 143857-10-7P

 RL: SPN (Synthetic preparation); PREP (Preparation) (preparation, crystal and mol. structure of)

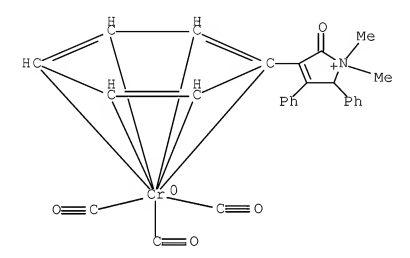
 RN 143857-10-7 HCAPLUS

CN Chromium(1+), tricarbonyl[2,5-dihydro-1,1-dimethyl-2-oxo-3-(η 6-phenyl)-4,5-diphenyl-1H-pyrrolium]-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 143857-09-4 CMF C27 H22 Cr N O4

CCI CCS



CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

CC 29-11 (Organometallic and Organometalloidal Compounds) Section cross-reference(s): 75 IT 57205-92-2P 136710-68-4P 138176-97-3P 138177-00-1P 143841-26-3P 143841-27-4P 143857-10-7P 143857-11-8P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation crystal and mol structure of)

(preparation, crystal and mol. structure of)

OSC.G 48 THERE ARE 48 CAPLUS RECORDS THAT CITE THIS RECORD (51 CITINGS)

L11 ANSWER 18 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1992:571276 HCAPLUS Full-text

DN 117:171276

OREF 117:29613a,29616a

TI Isoxazolidine compounds. XI. 2,2-Dialkyl-3-aryl-5-oxoisoxazolidinium salts: cyclic analogs of Polonovski intermediates

AU Steudle, Harald; Stamm, Helmut

CS Fak. Pharm., Univ. Heidelberg, Heidelberg, D-6900, Germany

SO Archiv der Pharmazie (Weinheim, Germany) (1992), 325(6), 329-31 CODEN: ARPMAS; ISSN: 0365-6233

DT Journal

LA German

OS CASREACT 117:171276

GΙ

AB Title salts I (R = Me, Et; R1 = H, Me; R2 = H, Me, CO2Me; X = H, Cl) were prepared in 45-95% yields by quaternization of the corresponding 2-methyl-3-aryl-5-oxoisoxazolidines with magic methyl or triethyloxonium tetrafluoroborate. The very sensitive cations I underwent solvolysis easily in MeOH without recognizable Polonovskii rearrangement; the products were 4-XC6H4CH:CR2CO2Me.

IT 143601-60-9P 143601-62-1P

143601-64-3P 143601-66-5P 143601-78-9P

143601-80-3P 143601-82-5P 143601-84-7P

143601-86-9P 143601-88-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and solvolysis of)

RN 143601-60-9 HCAPLUS

CN Isoxazolidinium, 2-ethyl-2-methyl-5-oxo-3-phenyl-, trans-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 143601-59-6 CMF C12 H16 N O2

Relative stereochemistry.

CM 2

CRN 14874-70-5

CMF B F4

CCI CCS

RN 143601-62-1 HCAPLUS

CN Isoxazolidinium, 3-(4-chlorophenyl)-2-ethyl-2-methyl-5-oxo-, trans-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 143601-61-0

CMF C12 H15 C1 N O2

Relative stereochemistry.

CRN 14874-70-5

CMF B F4

RN 143601-64-3 HCAPLUS

CN Isoxazolidinium, 2-ethyl-2,4,4-trimethyl-5-oxo-3-phenyl-, trans-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 143601-63-2

CMF C14 H20 N O2

Relative stereochemistry.

CRN 14874-70-5

CMF B F4 CCI CCS

RN 143601-66-5 HCAPLUS

CN Isoxazolidinium, 3-(4-chlorophenyl)-2-ethyl-2,4,4-trimethyl-5-oxo-, trans-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 143601-65-4

CMF C14 H19 C1 N O2

Relative stereochemistry.

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

RN 143601-78-9 HCAPLUS

CN Isoxazolidinium, 2-ethyl-1-methyl-5-oxo-3-phenyl-, cis-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 143601-77-8 CMF C12 H16 N O2

Relative stereochemistry.

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

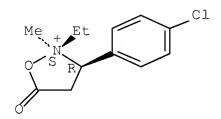
CN Isoxazolidinium, 3-(4-chlorophenyl)-2-ethyl-2-methyl-5-oxo-, cis-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 143601-79-0

CMF C12 H15 C1 N O2

Relative stereochemistry.



CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

RN 143601-82-5 HCAPLUS

CN Isoxazolidinium, 2-ethyl-2,4,4-trimethyl-5-oxo-3-phenyl-, cis-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 143601-81-4

CMF C14 H20 N O2

Relative stereochemistry.

CRN 14874-70-5

CMF B F4

RN 143601-84-7 HCAPLUS

CN Isoxazolidinium, 3-(4-chlorophenyl)-2-ethyl-2,4,4-trimethyl-5-oxo-, cis-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 143601-83-6

CMF C14 H19 C1 N O2

Relative stereochemistry.

CRN 14874-70-5

CMF B F4 CCI CCS

RN 143601-86-9 HCAPLUS

CN Isoxazolidinium, 2-ethyl-4-(methoxycarbonyl)-2-methyl-5-oxo-3-phenyl-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 143601-85-8 CMF C14 H18 N O4

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

RN 143601-88-1 HCAPLUS

CN Isoxazolidinium, 3-(4-chlorophenyl)-2-ethyl-4-(methoxycarbonyl)-2-methyl-5-oxo-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 143601-87-0 CMF C14 H17 C1 N O4

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

CC 28-6 (Heterocyclic Compounds (More Than One Hetero Atom))
IT 143601-60-9P 143601-62-1P

10/536,829

```
143601-64-3P
                    143601-66-5P
                                  143601-68-7P
     143601-70-1P 143601-72-3P 143601-74-5P 143601-76-7P
                   143601-80-3P 143601-82-5P
     143601-78-9P
     143601-84-7P
                   143601-86-9P 143601-88-1P
     143625-66-5P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
     RACT (Reactant or reagent)
        (preparation and solvolysis of)
    ANSWER 19 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN
     1992:428879 HCAPLUS Full-text
     117:28879
OREF 117:5199a,5202a
     Thermosetting resins containing cationic polymerization catalysts
     for coating materials
    Nakano, Shinji; Urano, Satoru; Osuqi, Koji
    Nippon Paint Co., Ltd., Japan
    Jpn. Kokai Tokkyo Koho, 17 pp.
    CODEN: JKXXAF
    Patent
    Japanese
FAN.CNT 1
    PATENT NO.
                        KIND
                               DATE
                                          APPLICATION NO.
                                                                  DATE
     _____
    JP 04023807
                        A 19920128 JP 1990-127782
ΡI
                                                                   199005
                                                                   17
                               19900517
PRAI JP 1990-127782
    MARPAT 117:28879
     The title polymerization catalysts are quaternary ammonium salts such
     as N-(\alpha, \alpha-dimethylbenzyl)pyridinium hexafluoroantimonate (I), N-(\alpha-
     methylbenzyl)-N,N-dimethylanilene hexafluoroantimonate, N-(\alpha-
     methylbenzyl)-pyridinium- hexafluoroantimonate, etc. Thus, 90 parts
     (solids) 1.88:2.59:28.11:25:30 iso-Bu methacrylate-Bu acrylate-Me
     methacrylate-styrene-glycidyl methacrylate copolymer was mixed with 2
     parts I, coated on tin plate, and baked to prepare a colorless
     coating.
     141915-34-6 141915-36-8
                                141915-38-0
     141915-41-5 141915-46-0
    RL: CAT (Catalyst use); USES (Uses)
        (catalysts, for thermosetting resins for coating materials)
    141915-34-6 HCAPLUS
     Oxazolium, 2,3-dihydro-3,3-dimethyl-2-phenyl-,
     (OC-6-11)-hexafluoroantimonate(1-) (9CI) (CA INDEX NAME)
```

CM 1

L11

ΑN DN

ΤI

ΙN PA

SO

DT

LA

OS

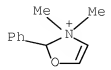
AΒ

ΙT

RN

CN

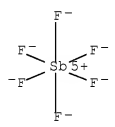
CRN 141915-33-5 CMF C11 H14 N O



CM 2

CRN 17111-95-4

CMF F6 Sb CCI CCS



RN 141915-36-8 HCAPLUS

CN Oxazolium, 2,3-dihydro-3,3-dimethyl-2-(4-methylphenyl)-, (OC-6-11)-hexafluoroantimonate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 141915-35-7

CMF C12 H16 N O

CRN 17111-95-4

CMF F6 Sb

RN 141915-38-0 HCAPLUS

CN Oxazolium, 3-ethyl-2,3-dihydro-3-methyl-2-(2-methylphenyl)-, (OC-6-11)-hexafluoroantimonate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 141915-37-9 CMF C13 H18 N O

CRN 17111-95-4

CMF F6 Sb CCI CCS

RN 141915-41-5 HCAPLUS

Oxazolium, 2,3-dihydro-2-(4-methoxyphenyl)-3,3-dimethyl-, CN (OC-6-11)-hexafluoroantimonate(1-) (9CI) (CA INDEX NAME)

CM 1

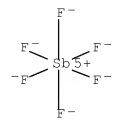
CRN 141915-40-4 CMF C12 H16 N O2

CM2

CRN 17111-95-4

CMF F6 Sb

CCI CCS

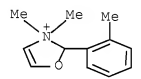


RN 141915-46-0 HCAPLUS

CN Oxazolium, 2,3-dihydro-3,3-dimethyl-2-(2-methylphenyl)-, hexafluorophosphate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 141915-45-9 CMF C12 H16 N O



CM 2

CRN 16919-18-9

CMF F6 P CCI CCS

IC ICM C08F004-00 ICS C08F004-06; C08G059-68

CC 42-10 (Coatings, Inks, and Related Products)

IT 136608-55-4 136608-64-5 136608-77-0 136842-16-5 141915-32-4 141915-34-6 141915-36-8 141915-38-0 141915-39-1 141915-41-5 141915-44-8

141915-46-0 141915-47-1 141915-49-3 141979-77-3

RL: CAT (Catalyst use); USES (Uses)

(catalysts, for thermosetting resins for coating materials)

L11 ANSWER 20 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1992:174051 HCAPLUS Full-text

DN 116:174051

OREF 116:29459a, 29462a

TI Thermolyses of 1,1-dimethyl-2-pyrazolinium fluoborates. Evidence for spiro[2,5]-1-aza-1,4,6-octatrienyl cation

AU Subramaniam, Girija

CS Dep. Chem., Penn State Univ., Hazleton, PA, 18201, USA

SO Indian Journal of Chemistry, Section B: Organic Chemistry Including Medicinal Chemistry (1992), 31B(3), 172-6 CODEN: IJSBDB; ISSN: 0376-4699

DT Journal

LA English

GΙ

AB Thermolysis of 3-aryl-1,1-dimethyl-2-pyrazolinium fluoroborates I (R = H, Me, MeO, Br, O2N) gives isomeric mixts. of 3-aryl-1-methylpyrazoles II and 5-aryl-1-methylpyrazoles III as major products

in complete contrast to the corresponding acyclic analogs. 2,6-Diaryl-3-methylpyridines IV were isolated only in trace quantities. The probable reasons for this unique behavior were explored using semi-empirical calcns., non-kinetic methods and radiolabeling expts. A pathway is proposed.

IT 139933-83-8

RL: RCT (Reactant); RACT (Reactant or reagent)
 (oxidation of)

RN 139933-83-8 HCAPLUS

CN 1H-Pyrazolium, 3-(4-aminophenyl)-4,5-dihydro-1,1-dimethyl-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 139933-82-7 CMF C11 H16 N3

$$Me^{N+1}$$

CM 2

CRN 14874-70-5

CMF B F4

IT 139933-70-3P 139933-72-5P

139933-74-7P 139933-76-9P 139933-78-1P

139933-85-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and thermolysis of, mechanism of) RN

139933-70-3 HCAPLUS

1H-Pyrazolium, 4,5-dihydro-1,1-dimethyl-3-phenyl-, CN tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM1

CRN 46186-33-8 CMF C11 H15 N2

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

RN 139933-72-5 HCAPLUS

1H-Pyrazolium, 4,5-dihydro-1,1-dimethyl-3-(4-methylphenyl)-, CN tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM1

CRN 139933-71-4 CMF C12 H17 N2

CRN 14874-70-5

CMF B F4
CCI CCS

RN 139933-74-7 HCAPLUS

CN 1H-Pyrazolium, 4,5-dihydro-3-(4-methoxyphenyl)-1,1-dimethyl-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 139933-73-6 CMF C12 H17 N2 O

CM 2

CRN 14874-70-5

CMF B F4

CCI CCS

RN 139933-76-9 HCAPLUS

CN 1H-Pyrazolium, 3-(4-bromophenyl)-4,5-dihydro-1,1-dimethyl-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 139933-75-8 CMF C11 H14 Br N2

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

$$-F - \frac{F}{B} - \frac{3}{3} + F$$

RN 139933-78-1 HCAPLUS

CN 1H-Pyrazolium, 4,5-dihydro-1,1-dimethyl-3-(4-nitrophenyl)-,

10/536,829

tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 139933-77-0 CMF C11 H14 N3 O2

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

RN 139933-85-0 HCAPLUS

CN 1H-Pyrazolium, 4,5-dihydro-1-methyl-1-(methyl-14C)-3-phenyl-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 139933-84-9

CMF C11 H15 N2

СМ

2

```
CRN 14874-70-5
     CMF B F4
     CCI CCS
CC
     28-8 (Heterocyclic Compounds (More Than One Hetero Atom))
     Section cross-reference(s): 22
     139933-83-8
ΙT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (oxidation of)
                    139933-72-5P
IΤ
     139933-70-3P
                    139933-76-9P 139933-78-1P
     139933-74-7P
     139933-85-0P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
     RACT (Reactant or reagent)
        (preparation and thermolysis of, mechanism of)
              THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1
OSC.G
              CITINGS)
L11
     ANSWER 21 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN
ΑN
     1991:429457 HCAPLUS Full-text
     115:29457
DN
OREF 115:5189a,5192a
     Reactions of fluorophosphoranes with
ΤI
     N, N, N'-trimethyl-N'-(trimethylsilyl)ethylenediamine.
     Intramolecularly stabilized azonium hexafluorophosphates by fluoride
     abstraction from N,N,N'-trimethylethylenediamine-substituted
     fluorophosphoranes with phosphorus pentafluoride
     Kaukorat, Thomas; Jones, Peter G.; Schmutzler, Reinhard
ΑU
     Inst. Anorg. Anal. Chem., Tech. Univ. Braunschweig, Braunschweig,
CS
     D-3300, Germany
SO
     Chemische Berichte (1991), 124(6), 1335-46
     CODEN: CHBEAM; ISSN: 0009-2940
```

DT Journal LA English

OS CASREACT 115:29457

GΙ

Reaction of the tetrafluorophosphoranes RPF4 (R = Me, Ph, C6F5, AB Me3SiCH2, 2,5-Me2C6H3) with N,N,N'-trimethyl-N'-(trimethylsilyl)ethylenediamine gave the corresponding trifluorophosphoranes I by cleavage of the Si-N bond and elimination of Me3SiF. In an analogous reaction the difluorophosphoranes R1R2PF2NMeCH2CH2NMe2 (R1 = R2 = Ph; R1 = Ph, R2 = C4H4N) are formed. Some of these N, N, N'-trimethylethylenediamine-substituted di- and trifluorophosphoranes react with PF5 as a Lewis acid to form the corresponding azonium hexafluorophosphates II (R = Me, Me3SiCH2, Ph, X = PF6) as a result of fluoride abstraction and intramol. Me2N \rightarrow P donor-acceptor interaction. II (R = NMe2, X = Cl) shows dynamic behavior in solution An exchange process is observed for the axial and equatorial fluorine atoms by 19F and 31P NMR spectroscopy. X-ray structure anal. of the compds. II reveals the expected trigonalbipyramidal geometry at phosphorus. The ethylenediamine ligand forms a chelate ring, whereby one axial and one equatorial site are bridged. The coordinate P-N bonds are very long (up to 207 pm).

IT 132750-93-7p 132750-95-9p

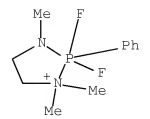
RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 132750-93-7 HCAPLUS

CN 1,3,2-Diazaphospholidinium, 2,2-difluoro-2,2-dihydro-1,1,3-trimethyl-2-phenyl-, stereoisomer, hexafluorophosphate(1-) (9CI) (CA INDEX NAME)

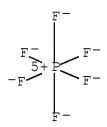
CM 1

CRN 132750-92-6 CMF C11 H18 F2 N2 P



CRN 16919-18-9

CMF F6 P CCI CCS



RN 132750-95-9 HCAPLUS

CN 1,3,2-Diazaphospholidinium, 2-fluoro-2,2-dihydro-1,1,3-trimethyl-2-phenyl-2-(1H-pyrrol-1-yl)-, stereoisomer, hexafluorophosphate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 132750-94-8 CMF C15 H22 F N3 P

CM 2

CRN 16919-18-9

CMF F6 P

CCI CCS

CC 29-7 (Organometallic and Organometalloidal Compounds)

Section cross-reference(s): 75

IT 19415-88-4P 132750-81-3P 132750-82-4P 132750-83-5P

132750-84-6P 132750-85-7P 132750-86-8P 132750-87-9P

132750-89-1P 132750-91-5P **132750-93-7**P

132750-95-9P 132750-96-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

OSC.G 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)

L11 ANSWER 22 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1988:186471 HCAPLUS Full-text

DN 108:186471

OREF 108:30627a,30630a

TI Synthesis and properties of cyclic keto alkenylammonium salts

AU Jung, Michael E.; Love, Brian E.

CS Dep. Chem. Biochem., Univ. California, Los Angeles, CA, 90024, USA

Journal of the Chemical Society, Chemical Communications (1987), (17), 1288-90

CODEN: JCCCAT; ISSN: 0022-4936

DT Journal

LA English

OS CASREACT 108:186471

GI

Condensation of RCH2COCHR1NMeR2 (R = H, Me, Et, Ph; R1 = H, Me, Ph; R2 = Me, Ph, CH2Ph) with HCO2Et in PhMe-NaH gave 33-100% sodium enolates I, which were cyclized by treatment with 4-MeC6H4SO2Cl in MeCN to give alkenylammonium salts II (R = H, Me, Et, Ph; R1 = H, Me, Ph; R2 = Me; Z= 4-MeC6H4SO3) in 41-88% yields. II [R = Ph, R1 = H, R2 = Me; R-R2 = Me; Z = MeSO3, (EtO)2PO2, BF4, ClO4] were prepared similarly.

IT 114050-22-5P

RN 114050-22-5 HCAPLUS

CN 3H-Pyrrolium, 1,2-dihydro-1,1-dimethyl-3-oxo-4-phenyl-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 114050-19-0 CMF C12 H14 N O

CM 2

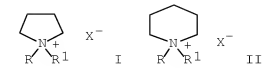
PRAI JP 1985-100929

GΙ

```
CRN 14874-70-5
    CMF B F4
    CCI CCS
CC
    27-10 (Heterocyclic Compounds (One Hetero Atom))
    114050-10-1P 114050-12-3P 114050-14-5P 114050-16-7P
ΙΤ
    114050-18-9P 114050-20-3P 114050-21-4P 114050-22-5P
    114050-23-6P 114050-24-7P 114050-25-8P
    RL: SPN (Synthetic preparation); PREP (Preparation)
       (preparation of)
OSC.G
            THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (5
            CITINGS)
L11 ANSWER 23 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN
AN
    1987:468042 HCAPLUS Full-text
    107:68042
DN
OREF 107:11081a,11084a
    Charge-donating materials for electrostatic image development
ΤI
IN
    Tanaka, Katsuhiko; Fukumoto, Hiroshi
PA Canon K. K., Japan
SO
    Jpn. Kokai Tokkyo Koho, 6 pp.
    CODEN: JKXXAF
DT
   Patent
LA
    Japanese
FAN.CNT 1
    PATENT NO. KIND DATE APPLICATION NO.
                                                      DATE
                                       ______
PI JP 61258269 A 19861115 JP 1985-100929
```

19850513

198505 13



The title materials contain a compound of the formula I or II (R, R1 = H, alkyl, aralkyl, aryl; X- = BF4-, PF6-) at least on their surfaces. The materials are useful for charging electrostatog. toners to provide adequate neg. charges. An Fe powder carrier was dispersed in a solution of II (R, R1 = tert-Bu; X = BF4-) (III) in MEK, dried, and the resulting coated carrier mixed with a toner prepared from D-125 (polystyrene) and Raven 3500 (C black) to give an electrostatog. developer, which had a triboelec. charge of -11 μ C/g as compared with -2 μ c/g for a control using a carrier containing no III. The developer gave high-quality images with good line reproduction and gradation and without fog and showed improved durability as compared with the control.

IT 109662-69-3

RL: USES (Uses)

(neq. charge-providing agent, for electrostatog. developers)

RN 109662-69-3 HCAPLUS

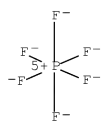
CN Pyrrolidinium, 1,1-diethyl-3-phenyl-, hexafluorophosphate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 109662-68-2 CMF C14 H22 N



CRN 16919-18-9 CMF F6 P CCI CCS



IC ICM G03G009-10 ICS G03G015-08

CC 74-3 (Radiation Chemistry, Photochemistry, and Photographic and Other Reprographic Processes)

IT 107751-93-9 107751-95-1 107751-96-2 107751-98-4 109662-64-8 109662-65-9 109662-67-1 109662-69-3

RL: USES (Uses)

(neg. charge-providing agent, for electrostatog. developers)

OSC.G 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

L11 ANSWER 24 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1984:423397 HCAPLUS Full-text

DN 101:23397

OREF 101:3709a,3712a

TI Synthesis and characterization of anhydro-1,1-dialkyl-5-hydroxy-3-phenoxy-1,2,4-triazolium hydroxides

AU Potts, Kevin T.; Kuehnling, William R.; Murphy, Peter

CS Dep. Chem., Rensselaer Polytech. Inst., Troy, NY, 12181, USA

SO Journal of Organic Chemistry (1984), 49(13), 2404-7 CODEN: JOCEAH; ISSN: 0022-3263

DT Journal

LA English

OS CASREACT 101:23397

GΙ

1,1-Dialkylhydrazines and PhOCCl:NCOCl react to give excellent yields of anhydro-1,1-dialkyl-5-oxo-3-phenoxy-1,2,4-triazolium hydroxides I (R = R1 = Me, Et, PhCH2; R = Me, R1 = Et). The reaction is regiospecific, and the same products are obtained from PhoC(:NH)NHNRR1 and phosgene. Thiophosgene and isocyanide dichlorides give exocyclic sulfur and nitrogen containing zwitterions, resp. Alkylation of the triazolium hydroxides occurs exclusively on N-2, and an O-Ph to N-Ph migration was observed at apprx.205°. Via dynamic NMR expts., the diasteriotopic methylene hydrogens of the benzyl groups attached to the quaternary nitrogen atom gave thermodn. exchange data of Ea = 21.7 \pm 0.7 kcal mol-1, ΔH .thermod. = 21.0 \pm 0.4 kcal mol-1, ΔS .thermod. = 6 \pm 1 eu, and ΔG .thermod. = 19.3 \pm kcal mol-1.

IT 90195-80-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

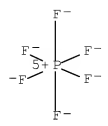
RN 90195-80-5 HCAPLUS

CN 3H-1,2,4-Triazolium, 1-ethyl-1,2-dihydro-2,2-dimethyl-3-oxo-5-phenoxy-, hexafluorophosphate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 90195-79-2 CMF C12 H16 N3 O2

CRN 16919-18-9 CMF F6 P CCI CCS



CC 28-9 (Heterocyclic Compounds (More Than One Hetero Atom)) Section cross-reference(s): 22

IT 90195-80-5P 90195-81-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

OSC.G 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)

L11 ANSWER 25 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1983:522390 HCAPLUS Full-text

DN 99:122390

OREF 99:18853a,18856a

TI Alkylation of the sulfur- and nitrogen-containing compounds analogous to thiazoline systems

AU Ohara, Yoshio; Akiba, Kinya; Inamoto, Naoki

CS Fac. Sci., Univ. Tokyo, Tokyo, 113, Japan

SO Bulletin of the Chemical Society of Japan (1983), 56(5), 1508-13 CODEN: BCSJA8; ISSN: 0009-2673

DT Journal

LA English

OS CASREACT 99:122390

AB RSCH2CH2NMe2 (R = alkyl) and 3-methylthiazolidines underwent N-methylation both with Me3O+BF4- and with MeI. With 2-MeSC6H4NMe2 the main reaction was S-methylation. On the other hand, in 2-EtSC6H4NMe2, S-methylated product was major with MeI, while N-methylated product was major with Me3O+BF4-. In the N-Et or N-benzyl derivative, only S-methylation occurred with both reagents. 3-Methyl-2,2-diphenyl-2,3-dihydrobenzothiazole gave S-methylated product with Me3O+BF4-, suggesting that S-methylation becomes major when the 2-substituent is bulkier.

4-Methyl-3,4-dihydro-2H-benzo[b][1,4]thiazine was N-methylated with Me30+BF4- while it gave S-methylated product with MeI. Phenothiazines and 1-dialkylamino-8-(methylthio)naphthalenes gave only S-methylated products. The possibility of intramol. interaction between sulfonio and amino groups has been discussed based on NMR data.

ΙT 74484-28-9P

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

74484-28-9 HCAPLUS RN

Thiazolidinium, 3,3-dimethyl-2-phenyl-, tetrafluoroborate(1-) (1:1) CN (CA INDEX NAME)

CM 1

CRN 74484-27-8 CMF C11 H16 N S

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

CC 28-14 (Heterocyclic Compounds (More Than One Hetero Atom))

Section cross-reference(s): 25

2525-18-0P 29898-80-4P 66623-87-8P 74484-24-5P 74484-26-7P ΙT

74484-28-9P 74484-30-3P 74484-32-5P 74484-36-9P

74484-41-6P 74484-39-2P 87094-23-3P 87094-25-5P 87094-26-6P 87094-27-7P 87094-29-9P 87094-31-3P 87094-32-4P 87094-33-5P 87094-35-7P 87094-36-8P 87094-38-0P 87094-39-1P 87094-40-4P 87094-43-7P 87094-45-9P 87094-42-6P 87094-47-1P 87094-49-3P 87094-51-7P 87094-52-8P 87094-53-9P 87094-55-1P 87094-56-2P 87094-60-8P 87094-58-4P 87094-62-0P 87094-66-4P 87111-91-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

OSC.G 6 THERE ARE 6 CAPLUS RECORDS THAT CITE THIS RECORD (6 CITINGS)

L11 ANSWER 26 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1982:216965 HCAPLUS Full-text

DN 96:216965

OREF 96:35833a,35836a

TI Two-step sigmatropic rearrangement versus aldol addition of ammonium imides

AU Gompper, Rudolf; Kohl, Bernhard

CS Inst. Org. Chem., Univ. Munich, Munich, D-8000/2, Fed. Rep. Ger.

SO Angewandte Chemie (1982), 94(3), 203-4 CODEN: ANCEAD; ISSN: 0044-8249

DT Journal

LA German

GI

H2NN+Me2CH2C(COPh):CHPh Br-, prepared from BrCH2C(COPh):CHPh and H2NNMe2, did not undergo a [2,3]-sigmatropic rearrangement, but rather formed I by intramol. aldol addition The intermediacy of the dipolar intermediate II in the [2,3]-sigmatropic rearrangement of HN-N+Me2CH2C(:CH2)COPh was demonstrated by the course of the reaction of PhCOC(CH2Cl):CH2 with Me2NNH2. Pyrazolidinium salt III was formed in Me2CO, but in Et2O salt IV was formed. Adding EtN(CHMe2)2 to the

ether gave 2:3 Me2NNHCH2C(:CH2)COPh(V)-IV. II was a common intermediate for IV and V; it was not completely certain that II was an intermediate for V.

IT 81917-60-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 81917-60-4 HCAPLUS

CN 1H-Pyrazolium, 4,5-dihydro-1,1-dimethyl-4-methylene-3-phenyl-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 81917-59-1 CMF C12 H15 N2

CM 2

CRN 14874-70-5

CMF B F4

CC 22-5 (Physical Organic Chemistry)

Section cross-reference(s): 28

IT 81917-57-9P **81917-60-4P** 81917-62-6P 81917-63-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

L11 ANSWER 27 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1982:35497 HCAPLUS Full-text

DN 96:35497

OREF 96:5881a,5884a

Organometallic chemistry of carbon-nitrogen multiple bonds. 3.

Reaction of tris(triphenylphosphine)platinum(0) with dimethylmethyleniminium chloride. X-ray structures of the products [(Ph3P)PtCH2N(CH3)2CH2N(CH3)2(Cl)]Cl and cis-[(Ph3P)Pt[CHN(CH3)2]Cl2]

AU Barefield, E. K.; Carrier, A. M.; Sepelak, D. J.; Van Derveer, D. G.

CS Sch. Chem., Georgia Inst. Technol., Atlanta, GA, 30332, USA

SO Organometallics (1982), 1(1), 103-10 CODEN: ORGND7; ISSN: 0276-7333

DT Journal

LA English

GΙ

Pt complex I was prepared in 78% yield by treating [Me2N:CH2]Cl with (Ph3P)3Pt in THF. Heating I in MeCN gave carbene cis-(Ph3P)Pt(:CHNMe2)Cl2 (II). The crystal structures of I and II were determined I has Pt-L distances of P, 2.223 (2); Cl, 2.362 (2); N, 2134 (4) and C, 2.017 (5) Å. The Pt-C distance is the shortest presently known for an sp3 C (without F substituents) bonded trans to Cl. II has Pt-L distances of P, 2.220 (2), Cl(1), 2.347 (3), Cl (2), 2.345 (3) and C, 1.96 (1) Å. The dihedral angle between the carbene and coordination planes is 84°.

IT 79328-84-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 79328-84-0 HCAPLUS

CN Platinum(1+), chloro[1-(dimethylamino)-N,N-dimethylmethanaminium methylide](triphenylphosphine)-, (SP-4-4)-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 79328-83-9

CMF C24 H31 Cl N2 P Pt

CCI CCS

$$\begin{array}{c} \text{Me} & \text{Me} \\ \text{H}_{2}^{-} \text{C} \\ \text{Ph}_{3} \text{P} & \text{-Cl} \\ \text{Me} \end{array}$$

CM 2

CRN 14874-70-5

CMF B F4

CCI CCS

CC 29-13 (Organometallic and Organometalloidal Compounds)

Section cross-reference(s): 75

IT 79328-84-0P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

OSC.G 20 THERE ARE 20 CAPLUS RECORDS THAT CITE THIS RECORD (20

CITINGS)

L11 ANSWER 28 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1981:515213 HCAPLUS Full-text

DN 95:115213

OREF 95:19325a,19328a

TI Reactions of pyryliums with mono- and asym-disubstituted hydrazines

AU Katritzky, Alan Roy; Ballesteros, Paloma; Tomas, Alberto Tarraga

CS Sch. Chem. Sci., Univ. East Anglia, Norwich, NR4 7TJ, UK

SO Journal of the Chemical Society, Perkin Transactions 1: Organic and

Bio-Organic Chemistry (1972-1999) (1981), (5), 1495-500

CODEN: JCPRB4; ISSN: 0300-922X

DT Journal LA English

OS CASREACT 95:115213

GΙ

The reactions of pyrylium salts with a variety of mono- and disubstituted hydrazines were studied. The products were generally either the appropriate pyridinium salt or the ring cleavage products; in 2 cases 2-pyrazoline derivs. were obtained. E.g., 2- ethoxycarbonyl-4,6-diphenylpyrylium tetrafluoroborate (I) with PhNHNH2 (MeOH, 20°, 30 min) gave 73% PhNHN:CPhCH2CPh:CHCOCO2Et, with H2NCONHNH2.HCl (aqueous EtOH, NaOH, reflux, 3 h) I gave the pyridinium salt II (56%), and with p-O2NC6H4NHNH2 (EtOH, 20°, 2 h) the product was III (59%). As was observed for several compds. derived from I, II readily cyclized [(MeOCH2)2, K2CO3, 20°, 2 h] to give 83% IV.

IT 78904-69-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 78904-69-5 HCAPLUS

CN 1H-Pyrazolium, 4,5-dihydro-1,1-dimethyl-5-(2-oxo-2-phenylethyl)-3,5-diphenyl-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 78904-68-4 CMF C25 H25 N2 O

CM 2

CRN 14874-70-5

CMF B F4

```
CC 27-17 (Heterocyclic Compounds (One Hetero Atom))
```

Section cross-reference(s): 23, 25, 28

IT 78904-65-1P 78904-66-2P 78904-67-3P 78904-69-5P

78904-70-8P 78904-71-9P 78904-73-1P 78904-75-3P 78904-76-4P 78904-77-5P 78904-78-6P 78904-79-7P 78904-80-0P 78904-81-1P

78904-82-2P 78904-84-4P 78904-90-2P 78904-92-4P 78904-97-9P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

OSC.G 7 THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD (7 CITINGS)

L11 ANSWER 29 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1980:495173 HCAPLUS Full-text

DN 93:95173

OREF 93:15257a,15260a

TI Alkylation of benzothiazolines and their Stevens rearrangement

AU Ohara, Yoshio; Akiba, Kinya; Inamoto, Naoki

CS Fac. Sci., Univ. Tokyo, Tokyo, 113, Japan

SO Fukusokan Kagaku Toronkai Koen Yoshishu, 12th (1979), 236-40 Publisher: Kitasato Daigaku Yakugakubu, Tokyo, Japan.

CODEN: 42VCA9

DT Conference

Japanese

LA GI

AΒ Benzothiazolines I (R = 4-MeOC6H4, 4-MeC6H4, Ph, 4-ClC6H4, Me, H) were methylated by Meerwein reagents to give benzothiazolinium salts Treating I with Et30+BF4- gave isomeric III. The selectivity of R30+BF4-, MeI, and MeI-AgClO4 in N- and/or S-alkylation of related compds. were examined Thus, when an N atom is in resonance with 2 benzene rings, i.e., phenothiazine, alkylation occurs on S even with R30+BF4-. Alkylation of N and S competes when the N is connected to Treating II with (Me2CH) 2NLi and BuLi caused a one C6H6 ring. Stevens rearrangement. Removal of a proton at C-2 by (Me2CH) 2NLi gave π -type cyclic ammonium ylidea as intermediates, which gave rearrangement products in moderate yield. Direct attack at the ring S by BuLi gave ammonium ylide IV, which gave mainly an unusual Stevens rearrangement to 2-(BuS)C6H4CHRNMe2 by participation of the ortho-BuS group.

IT 74484-28-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 74484-28-9 HCAPLUS

CN Thiazolidinium, 3,3-dimethyl-2-phenyl-, tetrafluoroborate(1-) (1:1) (CA INDEX NAME)

CM 1

CRN 74484-27-8

CMF C11 H16 N S

CM 2

CRN 14874-70-5

CMF B F4

```
28-7 (Heterocyclic Compounds (More Than One Hetero Atom))
CC
ΙT
    92-59-1P 614-30-2P 62290-08-8P 62290-10-2P
                                                   68549-59-7P
    70265-71-3P
                70265-72-4P 70265-73-5P 70459-24-4P
                                                        74484-13-2P
    74484-15-4P 74484-23-4P
                              74484-24-5P
                                           74484-25-6P
                                                        74484-26-7P
    74484-28-9P 74484-30-3P 74484-32-5P 74484-34-7P
    74484-36-9P 74484-39-2P
                              74484-41-6P 74484-42-7P 74484-43-8P
    74484-44-9P 74484-47-2P 74484-48-3P 74484-49-4P
                                                        74484-50-7P
                74484-52-9P 74484-54-1P
    74484-51-8P
    RL: SPN (Synthetic preparation); PREP (Preparation)
       (preparation of)
```

L11 ANSWER 30 OF 30 HCAPLUS COPYRIGHT 2009 ACS on STN

AN 1972:24570 HCAPLUS Full-text

DN 76:24570

OREF 76:3995a,3998a

TI Small charged rings. XV. Kinetics and stereochemistry of the ring expansion reaction of 2-arylaziridinium salts with benzaldehyde

AU Keenan, Thomas R.; Leonard, Nelson J.

CS Sch. Chem. Sci., Univ. Illinois, Urbana, IL, USA

SO Journal of the American Chemical Society (1971), 93(24), 6567-74

CODEN: JACSAT; ISSN: 0002-7863

DT Journal

LA English

OS CASREACT 76:24570

Preparation and isolation of 2-aryl-substituted aziridinium AB (ethyleniminium) salts are described for the first time. Ethyleniminium ions of this type were previously implicated as intermediates in adrenaline blockade. Using a series of 2-aryl-1,1dimethylaziridinium fluoroborates with BzH, the stereochemistry of the ring expansion reaction of aziridinium salts with aldehydes, was determined This representative of a family of aziridinium ring expansion reactions was highly stereoselective, since only his-5aryl-3,3-dimethyl-2-phenyloxazolidinium fluoroborates were produced. The kinetics of the reaction, determined by following the NMR spectra, showed the reaction to be first order in aziridinium salt, zero order in BzH, and dependent on the substituents on the aryl ring. A Hammett correlation with σ +, ρ + = -1.25 was observed stereochem. and kinetic findings preclude several mechanisms including concerted cycloaddn. and favor the intermediacy of aminocarbonium ions.

IT 34880-00-7P 34880-01-8P 34880-02-9P

34880-03-0P 34955-84-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 34880-00-7 HCAPLUS

CN Oxazolidinium, 3,3-dimethyl-2,5-diphenyl-, cis-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 46984-17-2 CMF C17 H20 N O

Relative stereochemistry.

CRN 14874-70-5

CMF B F4 CCI CCS

RN 34880-01-8 HCAPLUS

CN Oxazolidinium, 3,3-dimethyl-5-(4-methylphenyl)-2-phenyl-, cis-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 47068-18-8 CMF C18 H22 N O

Relative stereochemistry.

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

RN 34880-02-9 HCAPLUS

CN Oxazolidinium, 5-(4-chlorophenyl)-3,3-dimethyl-2-phenyl-, cis-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 47068-19-9

CMF C17 H19 C1 N O

Relative stereochemistry.

CM 2

CRN 14874-70-5

CMF B F4

CCI CCS

CN Oxazolidinium, 5-(4-bromophenyl)-3,3-dimethyl-2-phenyl-, cis-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 47068-17-7

CMF C17 H19 Br N O

Relative stereochemistry.

CM 2

CRN 14874-70-5

CMF B F4

CCI CCS

RN 34955-84-5 HCAPLUS

CN Oxazolidinium, 3,3-dimethyl-5-(4-nitrophenyl)-2-phenyl-, cis-, tetrafluoroborate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 47221-20-5

CMF C17 H19 N2 O3

Relative stereochemistry.

CM 2

CRN 14874-70-5

CMF B F4 CCI CCS

=>